

Empirical Threshold Values for Quantitative Trait Mapping

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ABSTRACT

The detection of genes that control quantitative characters is a problem of great interest to the genetic mapping community. Methods for locating these quantitative trait loci (QTL) relative to maps of genetic markers are now widely used. This paper addresses an issue common to all QTL mapping methods, that of determining an appropriate threshold value for declaring significant QTL effects. An empirical method is described, based on the concept of a permutation test, for estimating threshold values that are tailored to the experimental data at hand. The method is demonstrated using two real data sets derived from F2 and recombinant inbred plant populations. An example using simulated data from a backcross design illustrates the effect of marker density on threshold values.

INTRODUCTION

Methodological research on the problems of detecting and locating quantitative trait loci (QTL) has received considerable attention over the past several years. A variety of methods have been developed to analyze quantitative trait data (Weller 1986, 1987; Lander and Botstein 1989; Haley and Knott 1992; Knott and Haley 1992; Knott *et al.* 1994; Carbonell *et al.* 1992; Jansen 1993a,b; Jansen and Stam 1994; Zeng 1993, 1994). A problem common to all of these methods is the difficulty of determining appropriate significance thresholds (critical values) against which to compare test statistics (usually LOD scores or Likelihood Ratios) for the purpose of detecting QTL. The source of this difficulty is twofold. First, there is the problem of determining (or approximating) the distribution of the test statistic under an appropriate null hypothesis. In most cases, the regularity conditions that ensure an asymptotic chi-square distribution for the likelihood ratio test statistic are not satisfied. There are often additional problems due to finite sample sizes and distributional properties of the quantitative trait that might cause one to doubt the reliability of asymptotic approximations. The second source of difficulty is the multiple hypothesis testing that is implicit in the genome searches used for locating QTL. Large number of tests may be carried out, many of which are not independent. The dependence structure of these tests is difficult to analyze in cases other than the extremes of very dense or very sparse genetic maps. Elegant theoretical arguments have been presented (Lander and Botstein 1989) that address both of these issues. However they offer the user formulae for threshold values that are unfortunately difficult to apply and are based on a number of assumptions that are not likely to be met in practice.

The problem of determining appropriate threshold values is made even

more difficult because there are many factors that can vary from experiment to experiment and can influence the distribution of the test statistic. These include, but are not limited to, the sample size, the genome size of the organism under study, the genetic map density, segregation ratio distortions, the proportion and pattern of missing data and the distribution of the quantitative trait itself. Our goal in this work is to provide researchers involved in QTL mapping projects with a simple and intuitive procedure for estimating a threshold value and thus detecting significant QTL effects. Any such procedure must be statistically sound and should reflect, to the greatest extent possible, the characteristics of each particular experiment. In this paper, we describe a method based on the concept of permutation tests as first proposed by Fisher (1935). It involves repeated “shuffling” of the quantitative trait values and the generation of a random sample of the test statistic from an appropriate null distribution. The procedure is statistically valid when used in conjunction with likelihood or regression based test statistics and for *any* distribution of the quantitative trait. Because our procedure is empirical, based on the observed marker and trait data, it will automatically reflect the characteristics of the particular experiment to which it is applied.

Before proceeding to describe our method, we review the usual “QTL hypotheses” and discuss a handful of previous studies on the QTL detection problem, realizing that by no means are we presenting a complete literary synopsis of this field.

There are three hypotheses most relevant to the QTL detection problem (Knott and Haley 1992) these being

H_0^1 : no QTL is present.

H_0^2 : a QTL is present but is not linked to the marker(s) and

H_A : a QTL is present and is linked to the marker(s).

It is usual but not necessary to assume that, within classes defined by the (unknown) QTL genotype, the quantitative trait is normally distributed. Under the null hypothesis H_0^1 , the trait values should follow a single normal distribution. Thus any association between the trait values and a marker (interval) in the genetic map will be due purely to chance effects. Under the null hypothesis H_0^2 , the trait should follow a normal mixture distribution with mixing proportions equal to 1/2. Again any associations between the trait values and markers unlinked to the QTL are due to chance. Under the alternative hypothesis H_A , the trait should follow a normal mixture distribution with mixing proportions determined by the recombination fraction between the marker (interval) and the QTL. In this case, real associations between the trait values and the marker(s) are expected (Doerge 1993).

The most widely used algorithm for QTL detection and mapping is that implemented in the MAPMAKER/QTL software package (Paterson *et al.* 1988; Lincoln *et al.* 1992a,b) as first described by Lander and Botstein (1989). Their method is based on LOD scores (equivalent to log likelihood ratios) computed at regular incremental values throughout the genome. The null hypothesis assumed by MAPMAKER/QTL algorithm is H_0^1 . Although the null hypothesis of an unlinked QTL H_0^2 is discussed in the appendix A4 of Lander and Botstein (1989) as being more appropriate in some cases. An eloquent argument based on an Orenstein-Uhlenbeck diffusion process is used to determine the distribution of the maximum LOD score (over the entire genome) under the null hypothesis. Proposition 2 along with its Corrigendum (Lander and Botstein 1994) describes the calculation for determining the threshold value taking account of the known chromosome number and the known ge-

netic length of the organism. Lander and Botstein (1989) suggest that a *typical* LOD threshold should be between 2 and 3, to ensure an overall false positive (type I error) rate for QTL detection of 5%.

Knott and Haley (1992) used simulations to study the distributional properties of likelihood ratio tests for QTL detection. Their results suggest that the chi-square approximation to the distribution of likelihood ratio test statistics is not reliable in many cases and is at least questionable in every case. The problem of determining a significance threshold value for multiple nonindependent tests is not addressed in detail, but the reader is cautioned to consider setting higher significance thresholds in this case. In their conclusion Knott and Haley (1992) suggest that further theoretical work is needed in this area as no alternative other than simulation is presently available for setting significance thresholds.

Carbonell *et al.* (1992) note that it is inappropriate to use standard chi-square approximations for threshold values. They consider two different chi-square based thresholds and compare them to determine which gives “better” results. They also conclude that more research is needed to determine appropriate threshold values.

Zeng (1993) presents a regression based method that includes other markers as cofactors in a multiple regression. He advocates the use of an approximate one degree of freedom chi-square threshold for his method when the sample size is large, and the number of evenly spaced markers is small. Realizing that representative threshold values reflect the sample size, the number of markers in the model and the size of the marker interval, Zeng (1994) relies on a simulation study to explore these issues in a genome of unevenly markers and small sample size. For small sample size, the reader is cautioned as to the number of markers allowed in the model, since too many fitted markers can

substantially increase the threshold value of the test statistic. To this point Zeng offers no effective solution to the threshold problem other than to suggest using computer simulation.

Other researchers (Knapp *et al.* 1990; Jansen 1993b) have advocated the use of “conservative” threshold values based on chi-square distributions with either one or two degrees of freedom. The method presented by Jansen (1994) relies on weighted sum of squared residuals for the case of mixture models. Admittedly, Jansen (1994) states, “as an ad hoc approximation we used the chi-squared distribution with one degree of freedom, multiplied by the residual variance.” No justification is given for determining threshold values in this manner.

The difficulties in determining an appropriate asymptotic distribution are not too surprising when one considers that the most widely used hypothesis test compares a mixture distribution under H_A to a non-mixture distribution under H_0^1 . It is well documented (*e.g.* Hartigan 1985; Gosh and Sen 1985) that the usual asymptotic arguments do not apply in this case as the hypotheses are not properly nested. The fact remains that these hypotheses are of significant practical importance and thus we are highly motivated to study this problem further.

METHODS

Motivation: Data for quantitative trait analysis consist of a set of marker genotypes measured on each individual together with phenotypic trait values also measured on each individual. To detect QTL effects in the genome, statistical tests may be carried out at each marker and, if the markers are organized into a genetic map, at regularly spaced increments in the intervals between markers. We refer to the location at which a test statistic is computed as an

analysis point. In a *single marker analysis*, all of the analysis points are markers. If analysis points between markers are used, the analysis is an *interval analysis*. If there is a QTL effect at a specific location in the genome there will be an association between the trait values and the analysis points linked to that location. This association may be detected by t-tests or ANOVA (Scheffé 1959) in a single marker analysis or by likelihood ratios in an interval analysis. If there is no QTL effect linked to a marker, any associations between the trait values and the marker are likely to be weak and attributable to chance effects. Thus the key to detecting QTL effects is the detection of significant associations between the phenotypic trait values and the markers and/or intervals in a genetic map.

Our approach to the estimation of a significance threshold is based upon this simple observation of marker-phenotype association. It can be applied to single marker or interval mapping approaches using any test statistic with power to detect associations. If the data indicate that there are QTL effects, we can effectively destroy any association between the trait values and the analysis points linked to the QTL by randomly *shuffling* the trait values, *i.e.* by reassigning each trait value to a new individual while retaining the individual's genetic map. On the other hand, if the data fail to indicate the presence of QTL effects at specific regions of the genome, randomly shuffling the trait values across individuals will not alter the distribution of the test statistic. Any associations should still be small and attributable to chance. If we compute the value of an appropriate test statistic at each analysis point in the shuffled data sets, we are essentially sampling from a null distribution corresponding to the hypothesis of no associations between the trait values and the genetic maps. As the genetic maps and trait values are not altered by the shuffling procedure, this distribution will automatically take into account the particular

characteristics of the experiment at hand.

Threshold Estimation: Individuals in the experiment are indexed from 1 to n . The data are shuffled by computing a random permutation of the indices $1, \dots, n$ and assigning the i th trait value to the individual whose index is given by the i th element of the permutation. The shuffled data are then analyzed for QTL effects. The resulting test statistics at each analysis point are stored and the entire procedure (shuffling and analysis) is repeated N times. At the end of this process we will have stored the results of QTL analyses on N shuffled data sets. Two types of threshold values can be estimated from these results. The first is a comparisonwise threshold that can be estimated separately for each analysis point and provides a $100(1 - \alpha)\%$ critical value for the test *at that point*. The second is an experimentwise threshold that provides an overall $100(1 - \alpha)\%$ critical value that is valid simultaneously for all analysis points. Results of the QTL analysis on the original data can be compared to these critical values to determine statistical significance and thus to detect QTL effects.

A comparisonwise critical value is obtained by ordering the N test statistics obtained at each analysis point in the map and finding their $100(1 - \alpha)$ percentile. For example, if a comparisonwise significance level of $\alpha = 0.05$ is desired and $N = 1000$, the 950th value of the ordered test statistics will be our estimate of the comparisonwise critical value at that analysis point. Using this critical value to define a test controls the type I error rate *at that point* to be α or less. One should keep in mind that many individual tests may be computed and each presents a new opportunity to make a type I error. Thus if we use comparisonwise critical values, the type I error rate over the entire genome may be much higher than α .

The experimentwise critical value may be obtained by first finding the

maximum test statistic over all analysis points for each of the N shuffled analyses. These values are then ordered and their $100(1 - \alpha)$ percentile is our estimated experimentwise critical value. The experimentwise critical value is used to detect the presence of a QTL somewhere in the genome while controlling the overall type I error rate to be α or less. The experimentwise critical value will necessarily be higher than the comparisonwise values, thus the price for controlling the type I error rate over the entire genome is some loss of power to detect QTL effects.

An obvious question at this point is “How large should N be?”. We recognize that this procedure may be moderately expensive in computer time. Larger values of N will provide more precise estimates of the critical values. Thus there is a tradeoff here. Based on our limited experience with this procedure, we recommend that at least 1000 shuffles be used for estimating critical values at $\alpha = 0.05$. For more extreme critical values such as $\alpha = 0.01$, as many as 10,000 shuffles may be needed to obtain stable estimates.

Justification: The simple nature of the shuffling procedure seems almost too good to be true, but its use can be supported by well established statistical results. The procedure we have described is an approximate permutation test. Permutation tests were first proposed by Fisher (1935). A summary of the theory of permutation tests as well references to the original literature can be found in Lehmann (1986, pp. 230–245).

A permutation test in the simplest case is used to detect a location shift in data that are divided into two sets of observations $Z = (X_1, \dots, X_m, Y_1, \dots, Y_n)$. The test is performed by enumerating all permutations of the observed data $z' \in S(z)$ (where $S(z)$ denotes the set of all permutations of z) and ordering them by the values of a function $h(z')$ which is usually the likelihood of the data under an alternative hypothesis. The null hypothesis of no shift in

location is rejected if the value of the function $h(z)$ based on the observed data is among the k largest values of $h(z')$ based on the permuted data where $k = \lfloor \alpha n! \rfloor$, and $\lfloor x \rfloor$ denotes the greatest integer not greater than x . In practice the number of permutations is usually too large to enumerate. However, one can compute an approximate permutation test by generating a random sample from the set of all permutations of the data. For a random sample of size N , the estimated critical value is the k th largest value of $h(z')$ where $k = \lfloor \alpha N \rfloor$.

Consider testing for QTL effects at a single marker locus in a backcross population with a single segregating QTL. Let

$$Q_i = \begin{cases} 0 & \text{non-recurrent parental allele is absent at the QTL} \\ 1 & \text{non-recurrent parental allele is present at the QTL} \end{cases}$$

and

$$M_i = \begin{cases} 0 & \text{non-recurrent parental allele is absent at the marker} \\ 1 & \text{non-recurrent parental allele is present at the marker} \end{cases}$$

for individuals $i = 1, \dots, n$. Let Y_i be the phenotypic trait value of the i th individual. We will assume the trait value is a random variable with (conditional) density function $p_{Y|Q}(y, 0) = f(y)$ within the class of individuals defined by $Q_i = 0$ and with density function $p_{Y|Q}(y, 1) = f(y - \Delta)$ within the class of individuals defined by $Q_i = 1$. Thus the effect of the non-recurrent parental allele is to shift the density function of the trait by an amount Δ . We will assume, without loss of generality, that $\Delta > 0$. Of course, the QTL genotype cannot be directly observed. For a given marker linked to the QTL, with recombination fraction r ($0 \leq r \leq 1/2$), the conditional densities for the trait values are

$$p_{Y|M}(y, m) = r^m(1-r)^{1-m}f(y) + r^{1-m}(1-r)^mf(y-\Delta). \quad (1)$$

The mixture form of the density arises because the true QTL state is unknown. The essential points for our justification are

1. when $\Delta > 0$ and $r < 1/2$, $p_{Y|M}(y, 1)$ is stochastically larger than $p_{Y|M}(y, 0)$
2. when $\Delta = 0$ (corresponding to H_0^1) or when $r = 1/2$ (corresponding to H_0^2), the two conditional density functions, $p_{Y|M}(y, 0)$ and $p_{Y|M}(y, 1)$, are equal.

Now let

$$h(\mathbf{y}, \mathbf{m}) = \prod_{i=1}^n p_{Y|M}(y_i, m_i) \quad (2)$$

where $f()$ is taken to be a normal density function, $\mathbf{y} = y_1, \dots, y_n$ and $\mathbf{m} = m_1, \dots, m_n$. Thus $h()$ is the likelihood function under the alternative hypothesis H_A assuming the trait values are normally distributed within QTL genotype classes. Note that because $h()$ is used only to order the permutations, any monotone transformation of $h()$ such as the log likelihood, log likelihood ratio or LOD score will yield an equivalent test. With this choice of $h()$ and points (1) and (2) above, the conditions of lemma 3 in Lehmann (1986, p. 234) are satisfied. It follows that the permutation test is unbiased. That is, it has a type I error rate equal to α under either of the null hypotheses H_0^1 or H_0^2 and it has power greater than α for any alternative satisfying point (1) above. Furthermore, in the case where the true distribution of the phenotypic trait within the QTL genotype classes is normal, the permutation test is most powerful.

We note that the permutation test applied to a single marker is the non-parametric analog of a t-test. Although the t-test has proven to be robust, the conditions for a t-test are not satisfied in this case (Doerge 1993) and the

permutation test may be more appropriate. When the trait distribution is not normal within QTL genotype classes, the permutation test is still unbiased. In this case, a more powerful test could be derived by introducing the true density function $f()$ for the trait values into the equations (1) and (2). However, we note that the permutation test is robust to distributional assumptions and when applied with a normal density function $f()$ will generally lose very little power (Box and Anderson 1955).

This justification of the permutation test can be extended to analysis points within an interval. The mixtures in (1) are slightly more complex (Doerge 1993) but the stochastic ordering of the densities will still hold. In the case of F2 and other experimental crosses, the numbers of QTL genotype classes may be increased again making the mixtures in (1) more complex. The permutation test will still be unbiased for any form of the additive and dominance effects. However, because there are more than two marker classes, simple one-sided tests of location shift cannot be constructed and most powerful unbiased tests do not exist (Lehmann 1986). To apply the permutation test to the whole genome (experimentwise threshold), we consider a new function $h()$ which is the maximum of the likelihood over all analysis points. The conditions of lemma 3 (Lehmann 1986) are also satisfied by this function.

EXAMPLES

MAPMAKER Sample data: We have applied the permutation test to the F2 data that are distributed with MAPMAKER 1.1b software (Paterson *et al.* 1988; Lincoln *et al.* 1992a,b). The file `sample.raw` contains phenotypic trait data on 333 individuals and their genotypes at 12 marker loci. We used the linkage groups and map distances as established by the MAPMAKER/EXP 3.0 manual (Lander and Green 1987; Lincoln *et al.* 1992a,b).

The trait values were transformed by taking their logarithm for purposes of comparison with the sample analysis. (Transformation to obtain “normality” is not necessarily correct or even possible in some cases as the expected distribution of a trait in the presence of a QTL effect is a mixture distribution.) The trait values in `sample.raw` were shuffled and the usual QTL analysis was performed using MAPMAKER/QTL. This process was repeated 1000 times and the LOD scores at each 2cM increment analysis point were stored. To obtain comparisonwise threshold values, the 1000 LOD scores at each analysis point were sorted and the $100(1 - \alpha)$ percentile value was located. The results are summarized in table 1 and in figure 1. We see in this experiment that the comparisonwise threshold values are fairly constant across the two linkage groups. This need not be the case. Fluctuations in these values (shown in figure 1) are due in part to the sampling of the permutation set. They are larger for the more extreme critical values (e.g. $1 - \alpha = 0.99$) that are not as precisely estimated. Also note that the 95% LOD score threshold values are on average about 1.34. This LOD score can be rescaled to a likelihood ratio test statistic (divide by $\log_{10}(e)/2 = 0.2171$). The result 6.17 is slightly greater than the chi-square critical value on 2 degrees of freedom, $\chi^2_{2,0.95} = 5.99$. To obtain the experimentwise threshold, we first identify the maximum peak for each of the 1000 QTL analyses and then sort these values to obtain the $100(1 - \alpha)$ percentile (see table 1). The peak LOD score obtained on the original (not shuffled) data is 8.926, clearly indicating a significant QTL effect. A histogram of the 1000 maximum LOD scores is shown in figure 2. The distribution is seen to be right skewed as is typical for distributions of extreme values.

Single Marker Analysis of Recombinant Inbred Data: In this example, we consider a recombinant inbred (F7) population of rice derived from a cross between CO39 (maternal) and Moroberekan. A total of 203 recombi-

nant inbred lines were scored at 147 molecular markers. The quantitative trait of interest is root thickness (in microns). Details of the experimental methods can be found in Champoux, M.C., Wang, G., Sarkarung, S., Mackill, D.J., O'Toole, J.C., Huang, N., and McCouch, S.R. (unpublished).

Application of the permutation test to these data illustrates two points. First, it was noticed that the segregation ratios in this population are severely skewed and the experimentwise threshold values for QTL detection should reflect this peculiarity of the data. Second, interval mapping software for the analysis of RI populations is not readily available. Therefore, we have carried out a single marker analysis using a t-test at each analysis point (marker). We note that the assumptions of a t-test are not satisfied in this case. However the t-test is known to be robust to non-normality (Scheffé 1959; Doerge 1993) that arises due to the mixture distribution of the quantitative trait.

The original data were permuted 1000 times and the t statistics at each of the 147 markers were recorded. Comparisonwise thresholds were estimated for each of these tests (not shown). The degrees of freedom for each t-test vary slightly from marker to marker due to missing data. The averages of all 147 comparisonwise thresholds are summarized in table 2. Note that they compare very well with the corresponding t-distribution critical values. Experimentwise thresholds are of more interest in this example. The maximum t-test statistic (across all markers) from each of the 1000 permutations were used to obtain the experimentwise threshold. Results are summarized in table 2. The maximum t-test statistic for the original data was 9.0350, indicating a significant QTL effect in these data.

To determine if 1000 permutations of the data are sufficient to estimate experimentwise thresholds, we repeated the entire experiment 10 times. Standard errors of 0.028, 0.020 and 0.061 for the estimated threshold values were

obtained at $\alpha = 0.10, 0.05$ and 0.01 respectively. This suggests that 1000 permutations were adequate for estimating critical thresholds at $\alpha = 0.10$ and 0.05 . More extreme type I error rates such as $\alpha = 0.01$ may require larger numbers of permutations to yield threshold estimates accurate to two decimals.

A Simulated Example: One hundred backcross individuals were simulated in a genome containing four chromosomes of 100cM each. Chromosomes I and III were generated with 50 randomly placed markers each in a high density framework. Chromosomes II and IV were generated with a low density framework of 10 randomly placed markers each. The true genetic maps were used in subsequent QTL analysis. A QTL with additive effect 0.75 ($\sigma^2 = 1.0$) was simulated at 44.4cM from the left end of chromosome I. A second QTL of effect 1.0 ($\sigma^2 = 1.0$) was simulated at 61.6cM from the left end of chromosome II.

Each of the 1000 permuted data sets was analyzed in MAPMAKER/QTL under the backcross data type. Comparisonwise thresholds for the LOD scores are shown in figure 3 and the average values are summarized in table 3. Note that the comparisonwise values are fairly constant throughout the entire genome. To illustrate the effect of marker density on experimentwise thresholds, we have computed separate thresholds for each linkage group. These “linkage group-wise” thresholds correspond to four independent tests of QTL effects on each chromosome (and thus four opportunities to make a type I error). They are shown in figure 3 and summarized in table 3. For linkage groups II and IV with low density marker frameworks, the threshold values are lower. This point illustrates that the estimated threshold values do indeed reflect the characteristics of the experiment (chromosome) to which they are applied. We note also that the thresholds for chromosomes I and II that contain QTL

are essentially identical to thresholds for chromosomes III and IV that do not contain QTL. This last point illustrates that the shuffling effectively breaks up any association of QTL effects and markers.

DISCUSSION

We have presented a method of estimating threshold values for declaring significant QTL effects in a genome or at any point within a genome by the application of a permutation test. The test is valid for any continuously distributed trait, *i.e.* it will have the correct type I error level and will have power to detect QTL effects under the alternative hypothesis H_A . The threshold values obtained through this method are limited specifically to the data set on which the permutations are performed. However, since the method can be automated, one can easily generate threshold values for any data set. As an example of the time required for computing threshold values, our first example took approximately 23 hours of CPU time to generate 1000 QTL analyses of permuted data on a SPARC IPX Workstation. This is a relatively small investment compared to the many hours required to score and genotype individuals in a typical QTL experiment.

Generalizations of the permutation test to the problem of detecting multiple QTL effects may be possible. Lehmann (1986) notes that in the case of a heterogeneous population, the power of a permutation test can often be improved by stratifying the population according to factors unrelated to the treatment of interest but known to affect the outcome (analogous to unlinked QTL). In the case of QTL detection, the presence of a major gene(s) affecting the trait of interest may be known *a priori*. The population can then be divided into classes based on the presence/absence of the major gene(s) (or tightly linked marker) and the trait values permuted within these classes. This

procedure could significantly increase the power for detecting unlinked QTL effects secondary to the major gene. We caution potential users of this approach that if the classes are determined in light of the data from the present experiment, the type I error level of the procedure cannot be guaranteed. Also, one should avoid conditioning on too many markers in the genome. As the level of stratification increases or if the stratification is not effective in reducing the variance of the trait, the loss of power due to stratification may offset any advantages of conditioning.

The problem of detecting and locating multiple QTL effects has been most recently addressed by Jansen (1993a,b, 1994), Haley *et al.* (1994), and Zeng (1993, 1994). Each of these works makes us increasingly aware of the growing importance of determining threshold values against which to compare test statistics. Jansen (1994) states with regard to an overall significance value, “Many tests are performed when moving along the genetic map. An overall significance level cannot be guaranteed due to the lack of knowledge about the statistical behavior of the (interdependent) tests.” Similarly, Haley *et al.* (1994) suggest probing the null hypothesis distribution using Monte Carlo simulation for the purpose of getting at the distribution of their test statistic under the null hypothesis within the setting of multiple correlated test. At first glance, it appears that Haley *et al.* might be doing a permutation test, but this is not the case. At each Monte Carlo simulation, they are generating new phenotypic data (not permuting). Indeed, this suggestion will supply threshold values, but since the phenotypic data are being constructed, the particularities of the experimental situation such as segregation distortion and missing data may be lost and the resulting thresholds are highly model dependent. Lastly, Zeng (1993) asks in reference to his multiple regression method using cofactors, “since it is a multiple test and search problem (for multiple locations), what

would be an appropriate significance value of the test statistic given a data set?”

Clearly further work on obtaining threshold values is needed in order to answer these concerns. The permutation test offers the mapping community an intuitive method for estimating threshold values which accurately reflect the specifics of an experimental situation. The power of the permutation test is optimized when the function $h()$ is the true likelihood function for the data. It will be of some interest to compare the use of normal mixture likelihoods to other mixture distributions in this context. Further work is needed on the problems of modeling QTL effects, especially with regard to the multiple QTL detection problem. The reader will note that important problems of locating QTL and of estimating model parameters are not addressed in the present work.

In summary, the permutation test provides an easy to use method for estimating threshold values that is statistically sound, robust to departures from standard assumptions and is tailored to the experiment at hand. While the permutation test has been presented here in conjunction with MAPMAKER/QTL, it is completely feasible to use the permutation test with any QTL mapping procedure including simple linear regression, multiple regression and multiple regression with cofactors.

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Table 1: Estimated Threshold Values^a for MAPMAKER/QTL Sample Data

$1 - \alpha$	Experimentwise	Comparisonwise ^b
0.90	2.07	1.01
0.95	2.51	1.34
0.99	3.24	2.13

^aBased on LOD scores from 1000 permutations of the original data.

^bAverage across all analysis points.

Table 2: Estimated Threshold Values^a for Root Thickness Data

$1 - \alpha$	Experimentwise	Comparisonwise ^b
0.90	3.40	1.65
0.95	3.60	1.98
0.99	3.99	2.60

^aBased on t-statistics from 1000 permutations of the original data.

^bAverage across all analysis points.

Table 3: Estimated Threshold Values^a for the Simulated Backcross Data

$1 - \alpha$	Experimentwise				Comparisonwise ^b			
	LG1 ^c	LG2	LG3	LG4	LG1	LG2	LG3	LG4
0.90	1.45	1.29	1.48	1.31	0.58	0.61	0.59	0.61
0.95	1.83	1.57	1.84	1.59	0.84	0.87	0.85	0.86
0.99	2.44	2.21	2.51	2.19	1.48	1.43	1.51	1.45

^aBased on LOD scores from 1000 permutations of the original data.

^bAverage across all analysis points in the linkage group.

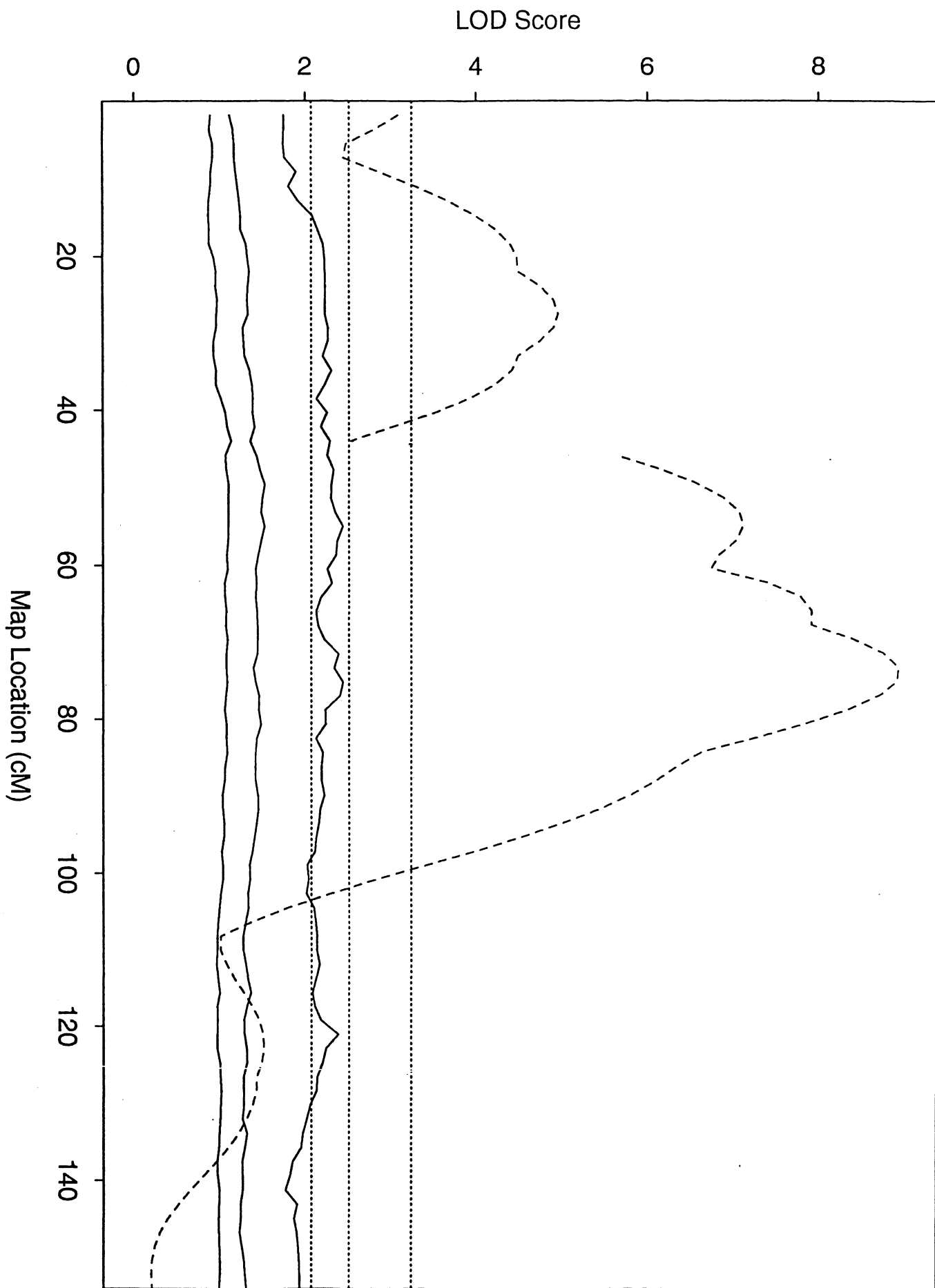
^cLG = Linkage Group.

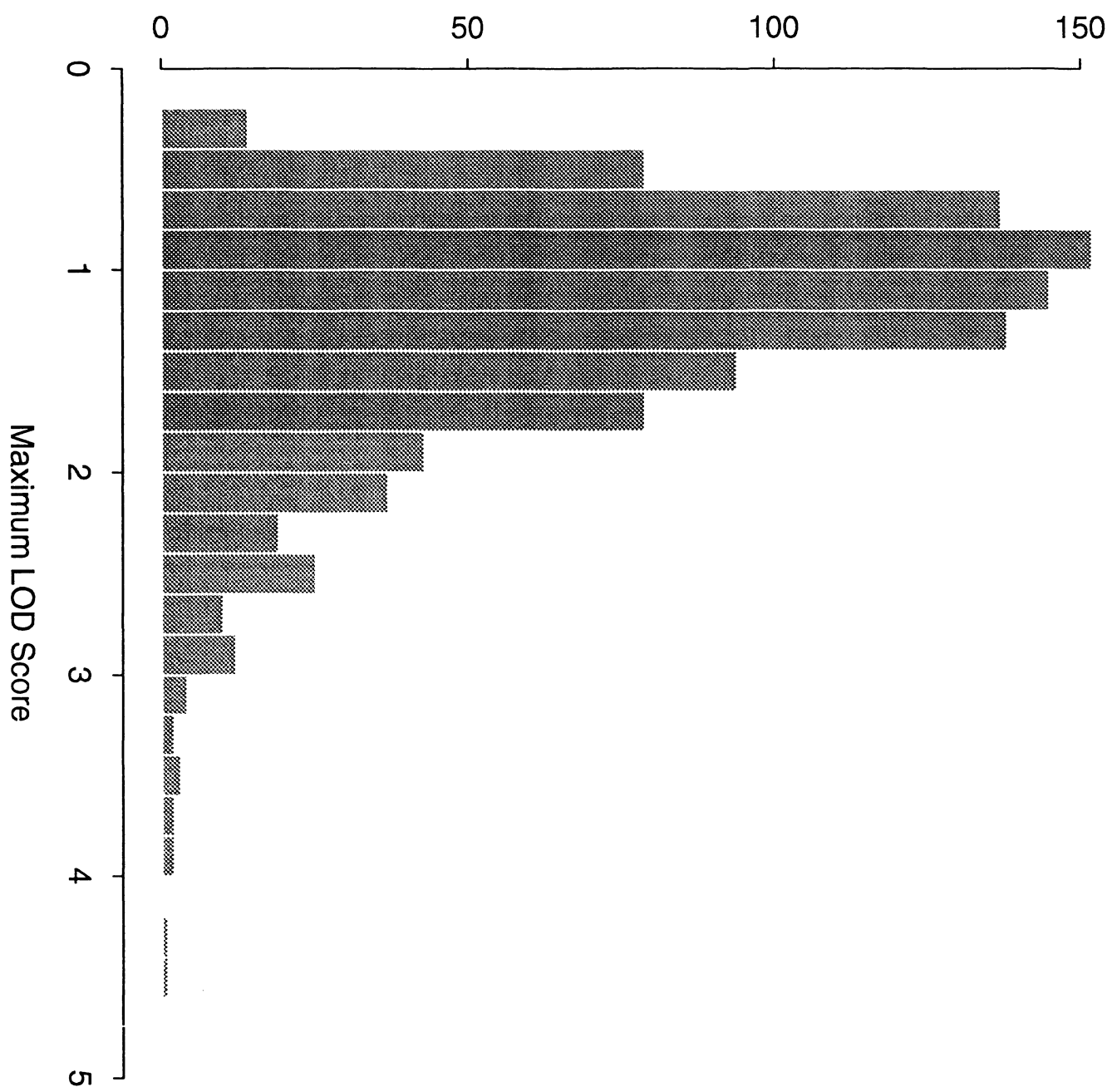
Figure Captions

Figure 1: QTL analysis of MAPMAKER/QTL sample data. LOD scores from analysis of the original data (\cdots) are plotted at each analysis point. Comparisonwise ($—$) and experimentwise ($- - -$) threshold values are shown for level $\alpha = 0.01$ (top), 0.05 (middle) and 0.10 (bottom) tests.

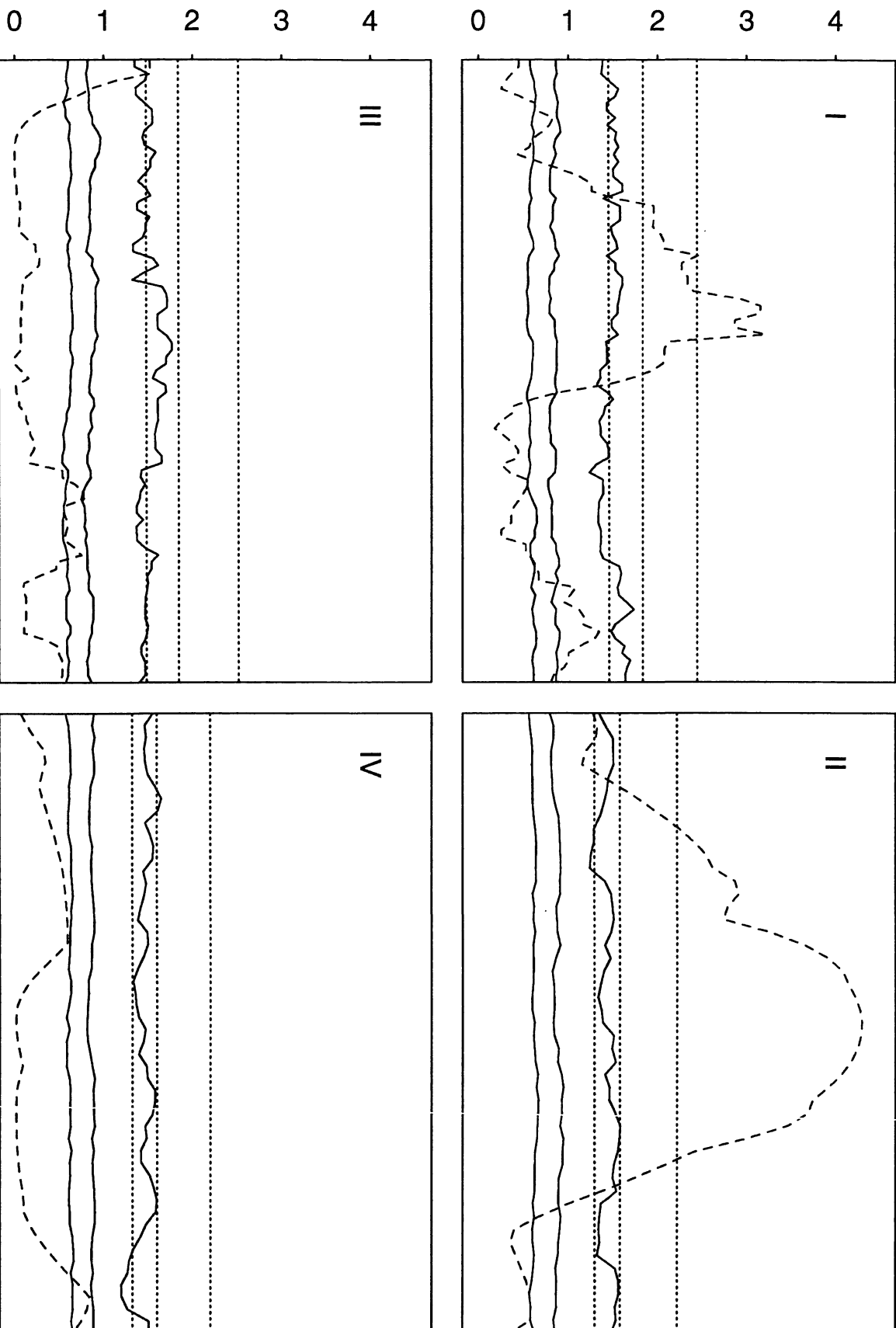
Figure 2: Histogram of maximum LOD scores for MAPMAKER/QTL sample data. The maximum LOD score across all analysis points was computed for each of 1000 permutations of the original data. Percentiles of this distribution are used to define experimentwise threshold values.

Figure 3: QTL analysis of simulated backcross data. The figure is divided into four sections corresponding to the four linkage groups. Each linkage group is 100cM in length. Linkage groups I and II contain QTLs. Linkage groups III and IV do not. Linkage groups I and III are mapped at high density (50 markers each) and linkage groups II and IV are mapped at low density (10 markers each). LOD scores from analysis of the original data (\cdots) are plotted at each analysis point. Comparisonwise ($—$) and linkage group-wise ($- - -$) threshold values are shown for level $\alpha = 0.01$ (top), 0.05 (middle) and 0.10 (bottom) tests.





LOD Score



Map Location (cM)